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Re-submitted to *Journal of Molecular Evolution**Research article*Tandem duplications in the C-terminal domain of the mesotocin receptor exclusively
identified among East Eurasian thrushesHideaki Abe^{1, 3}, Isao Nishiumi² and Miho Inoue-Murayama^{1, *}¹*Wildlife Research Center, Kyoto University, Kyoto, 606-8203, Japan*²*Department of Zoology, National Museum of Nature and Science, Tsukuba, 305-0005,
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Abstract

Mesotocin is a neurohypophyseal hormone found in some non-mammalian vertebrates, including birds, reptiles and amphibians. In this study, we identified and characterized 18-amino acid duplications in the C-terminal domain of the mesotocin receptor (MTR), specifically found in *Turdus* thrushes (Aves: Passeriforms: Turdidae). These duplicated elements are located in the distal part of the C-terminal tails of MTR and consist of amino acids that are highly conserved among major vertebrates. Intraspecific polymorphisms in a variable number of tandem duplications are commonly found in East Eurasian *Turdus*, but not in any other genus of Turdidae. Moreover, the genus *Turdus* can be further classified into 2 groups according to the presence or absence of a 3-amino acid deletion just adjacent to the putative palmitoylation site in the cytoplasmic C-terminal tail. The phylogeny presented here strongly supports the conspecific group of 4 East Eurasian thrushes (*Turdus pallidus*, *T. chrysolaus*, *T. obscurus*, and *T. celanops*). Our findings, therefore, provide a new synapomorphy that can be used for phylogenetic assumptions and shed a light on the history of diversification within Eurasian *Turdus* clades.

Keywords: mesotocin receptor — C-terminal domain — tandem duplication — polymorphism — insertions and deletions — *Turdus*

54 Introduction

55

56 Mesotocin (MT) is the non-mammalian homolog of mammalian oxytocin, and this neurohypophysial

57 hormone exerts biological effects by binding to G protein-coupled receptors (GPCRs) on the cell

58 surface. GPCRs are structurally characterized by 7 transmembrane domains, intracellular and

59 extracellular loops, and amino- and C-terminal segments (Gimpl and Fahrenholz 2001; Lolait et al.

60 1992). We focused on length variations in the C-terminal domain of peptide hormone receptor as a

61 key factor in influencing signal transduction pathways. Hoare et al. (1999) experimentally illustrated

62 that artificial truncation in the C-terminal tail of the human oxytocin receptor has critical effects in

63 receptor-G protein interactions. Furthermore, our recent study demonstrated that the arginine

64 vasotocin receptor has inter- and intraspecific variations in the length of C-terminal domains,

65 suggesting conformational differences in this region (Abe et al. 2012). Thus, we consider it worth

66 investigating whether the mesotocin receptor (MTR) also has conserved sequence motifs and amino

67 acid residues in the cytoplasmic C-tail.

68 In addition to characterization of novel polymorphisms, in this study we also present a hypothesis of

69 phylogenetic relationships among East Eurasian thrushes by taking advantage of synapomorphic

70 feature of insertions and deletions (indels) that are specifically found in *Turdus* clades. A

71 synapomorphic character is one of the most reliable piece of information on phylogenetic

72 reconstitution, as long as parallel changes and/or reversions are unlikely between different taxa. Not
73 only morphological characteristics but also genomic changes can be used as a landmark for
74 identifying a taxonomic clade. Rare genomic changes (RGCs) occasionally provide unambiguous
75 evidences for resolving the puzzle of phylogenetic problems (reviewed in Boore 2006; Rokas and
76 Holland 2000). RGCs can be caused by various molecular events—retroposon integrations, gene
77 order changes, and large-scale indels, etc. Some retroposons known as long and short interspersed
78 repetitive elements (LINEs and SINEs, respectively) have attracted special attention of evolutionary
79 biologists, because copies of these molecules shared at the same locus in 2 or more different taxa are
80 derived from the same element originally inserted into the germline of a common ancestor (Shedlock
81 and Okada 2000). Indeed, the presence or absence of SINEs insertions have been used for
82 phylogenetic analyses in a variety of organisms such as salmon (Kido et al. 1991), cetaceans
83 (Nikaido et al. 1999), and avian species (St. John et al. 2005; Watanabe et al. 2006). Moreover, some
84 amino acid indels and substitutions that had occurred in functional proteins proved to be
85 synapomorphic characters for avian clades (Ericson et al. 2000; Groth and Barrowclough 1999;
86 Stapel et al. 1984).

87 The genus *Turdus* is the largest group of Turdidae and perhaps the most successful of any passerine
88 genus in the world in terms of natural geographical penetration and degree of speciation. *Turdus* is
89 the only thrush genus that is widely distributed throughout Eurasia, South and North America, and

90 Africa. Of its nearly 70 species, 8 are confined to Africa, 25 are essentially Asian, and 34 are
91 restricted to the America (Collar 2005). A previous study on phylogenetic relationships among true
92 thrushes indicated that *Turdus* radiations occurred near the end of the Miocene (7–8 million years
93 ago) and rapidly diversified into each continental lineage (Klicka et al. 2005). Although the
94 subsequent statistical analysis based on mitochondrial genes indicated a monophyly of each
95 continental clade, relatively weak nodal support was found within the Eurasian *Turdus* clade
96 (Voelker et al. 2007). Such a rapid radiation and speciation, which are general characteristics of
97 avian evolution, always hamper phylogenetic approaches using stochastic methods (see Poe and
98 Chubb 2004). Another striking feature of the genus *Turdus* is its apparent plumage
99 homogeneity—similar plumage patterns appear repeatedly across its wide range. This makes it more
100 complicated to define species boundaries and unravel the evolutionary history. Therefore, it is
101 important to gather information on RGCs, because one can obtain invaluable data sets for cladistic
102 assumptions if synapomorphic characters could be found elsewhere within problematic taxa. This
103 approach is expected to provide a better resolution of phylogenetic relationships and interesting
104 insights into speciation processes and timing at multiple taxonomic levels.

105

106

107 Material and methods

108

109 *Avian specimens and DNA extraction*

110

111 A wide range of avian species were used in this study to explore sequence and structural features of
112 the C-terminal tails of the MTR. In addition, more than 200 *Turdus* specimens were particularly used
113 to further investigate intraspecific polymorphisms in this domain of the receptor. These specimens
114 were divided into 3 categories, depending on their sources: (1) live resources derived from zoos and
115 aviaries, (2) carcasses originated either from wildlife conservation acts or from bird-strike events in
116 Japanese airports, and (3) specimens obtained from avian research projects conducted by the
117 National Museum of Nature and Science. Supplementary S-Table 1 summarizes the avian samples
118 used for sequence alignment and their GenBank IDs. When we collected specimens from live
119 animals, we strictly observed the domestic laws for animal rights and protection in Japan. DNA was
120 extracted from various materials such as feathers, blood, and tissues (e.g., muscle or liver) using
121 either of the 3 methods: DNeasy Blood and Tissue Kit (Qiagen, Tokyo, Japan), Puregene DNA
122 Purification Kit (Gentra Systems, Minneapolis, MN, USA), or standard phenol-chloroform
123 extraction (Sambrook et al. 1989).

124

125 *Polymerase chain reaction amplification*

126

127 To amplify the C-terminal region of the MTR, primers were designed on the basis of the nucleotide
128 sequences of the zebra finch deposited in GenBank (Accession No. XM_00218826). The following
129 oligo primers were designed for amplification of avian *MTR*: MTf2 (5'–
130 GCCTCCCCCTTCATCATCG–3'), MTr3 (5'–AAGCTCCTGTGGCTCGTG–3'), MTr4 (5'–
131 CGGCAGCTGAGCACGAAG–3'), and MTr7 (5'–TCGCGGGCGGCTGCACGAA–3'). We mainly
132 used the MTf2/MTr3 primer pair, but MTf2/MTr4 was employed for amplification of Phasianidae
133 (Galliformes) and Anatidae (Anseriformes). MTf2/MTr7 was used in the amplifications of *Turdus*
134 thrushes, because the MTr3 annealing site overlapped with lineage-specific tandem duplicated
135 elements. In each case, polymerase chain reaction (PCR) was carried out in a PCR cocktail
136 containing approximately 10–50 ng of genomic DNA, 0.25 μ M each primer, 0.2 mM dNTPs, 5 \times
137 tuning buffer, and 0.375 U of G-Taq DNA polymerase (Hokkaido System Science, Hokkaido, Japan).
138 PCR conditions comprised an initial incubation at 95°C for 2 min, followed by 40 cycles of 95°C for
139 15 s and 60°C for 30 s.

140

141 *Sequencing and multiple alignments*

142

143 The nucleotide sequences of the C-terminal domain of MTR were determined in a total of 44 avian

144 species belonging to 31 families (17 orders). Sequencing was performed on an ABI 3130xl
145 automated sequencer (Applied Biosystems, Foster City, CA, USA) using BigDye Terminator Cycle
146 Sequencing reagents v3.1, according to the manufacturer's instructions (Applied Biosystems).
147 Nucleotide sequences were translated into amino acids and then aligned by using CLUSTAL W
148 (Thompson et al. 1994) implemented in BioEdit ver. 7.9.9.0 (Hall 1999). Each band of the amplified
149 alleles was purified from the gel and subjected to DNA sequencing. Information on amino acid
150 sequences of the C-terminal tail of vertebrate oxytocin and MT receptors were collected either from
151 previous papers (Akhundova et al. 1996; Hoare et al. 1999) or from GenBank sequences.

152

153 *Genotyping and sex determination*

154

155 Multiple sequence alignment revealed the existence of tandem duplicated elements in the distal
156 C-terminal tails of MTR specifically in *Turdus* thrushes. Thus, the pattern of length polymorphisms
157 was further investigated by genotype analysis using capillary electrophoresis. MTf2 was
158 fluorescently labeled with 6-carboxyfluorescein (6-FAM), and the HD500 Rox size standard
159 (Applied Biosystems) was used as an internal standard. PCR products were electrophoresed on an
160 ABI 3130xl sequencer, followed by peak detection with the Genemapper software (Applied
161 Biosystems). When we could not judge the gender according to the external features of the cloaca,

sexes were determined using the DNA-based identification method described by Griffiths et al. (1998). The exact fragment size of partial CHD-W and CHD-Z on automated capillary electrophoresis was taken from the empirical data of Lee et al. (2010). We evaluated the extent of statistical heterogeneity in allelic distributions between sampling locations (west Honsyu [$n = 39$] vs. Kyusyu [$n = 14$]) and sexes (male [$n = 55$] vs. female [$n = 29$]) of the pale thrush (*Turdus pallidus*) using Fisher's exact test.

Results

Structural characteristics in the C-terminal domain of the avian mesotocin receptor

In this study we determined the nucleotide sequences of the C-terminal domain of MTR in 44 avian species belonging to 17 orders. These sequences were deposited in the GenBank database (Accession Nos. AB634778–AB634824, and AB743549). We did not find any stop codon or frame-shift mutation in the sequence alignment; hence, we excluded the possibility that these sequences could be derived from MTR pseudogenes. The schematic secondary structure of the target region is shown in Figure 1. Multiple alignments showed that the distal part of TM VII (position

180 318–335) and its juxtamembrane region (position 336–343) are completely conserved among all
181 avian species (Fig. 2). The conserved NPxxY motif, which is characteristic of rhodopsin GPCRs
182 (Fritze et al. 2003), could be identified in TM VII (position 328–332). A double cysteine motif in the
183 C-terminus (positions 349, 350) was conserved in all of the avian and mammalian species; these
184 residues have been suggested by the CSS-Palm software (Ren et al. 2008) to be putative sites for
185 palmitoylation. Passerine birds could be distinguished from other avian species because of the
186 common deletion of 2 amino acids in the middle of the C-terminal domain (positions 364, 365).
187 Furthermore, we found 54-bp duplicated elements (18 amino acids; KSNSSSFVLSCRSPSHRS) in
188 *Turdus* thrushes. There were 4 types of alleles with the tandem repeat number varying from 1 to 4
189 [*NR* (wild-type), *RT2*, *RT3*, and *RT4*]. Four alleles were different by exactly 60-bp intervals. Global
190 alignment with other vertebrates uncovered that the duplicated region was highly conserved not only
191 among avian species but also among the mammalian species (Fig. 3). In particular, the
192 KSXSSXFVLS (X: arbitrary amino acid) motif was completely conserved in all vertebrates except
193 teleost fish. Although the functional significance of repeated motifs remains speculative, it should be
194 mentioned that tandem duplicated motifs contain consensus sites for phosphorylation by the
195 multifunctional protein kinase II (RxxS; Kemp and Pearson 1990).
196
197 *Distribution of tandem duplicated alleles within Turdus thrushes*

198

199 We detected intraspecific variations in the number of tandem duplications among *Turdus* thrushes,
200 whereas the White's thrush (*Zoothera dauma*) was monomorphic with the wild-type (i.e.,
201 non-duplicated) allele (Table 1). The other following avian species tested for length polymorphisms
202 did not show any variation in this locus (zebra finch [*Taeniopygia guttata*; $n = 32$], Bengalese finch
203 [*Lonchura striata* var. *domestica*; $n = 15$], red-throated parrot-finch [*Erythrura psittacea*; $n = 8$],
204 canary [*Serinus canaria*; $n = 10$], rock eagle-owl [*Bubo bengalensis*; $n = 8$], and mallard [*Anas*
205 *platyrhynchos*; $n = 8$]). These data indicate that duplication events occurred in the common ancestor
206 of *Turdus* thrushes. There was little difference in the allelic frequency among Eurasian *Turdus*
207 species. *RT2* was the predominant allele with a frequency ranging from 0.62 to 0.74; the frequency
208 of the other alleles was similar among species.

209

210 *Phylogenetic assumption based on rare genomic changes*

211

212 In addition to tandem duplications in the C-terminal tails, we found a 3-amino acid deletion (Ser-
213 Thr-Arg) at residues 351 to 353, just adjacent to a putative palmitoylation site (see Figs. 1 and 2).
214 The non-duplicated and duplicated alleles with this deletion were expressed as *NR-*, *RT2-*, *RT3-*,
215 and *RT4-*, and the frequency of each allele is shown in Table 1. The deletion was conserved in all

216 individuals belonging to 4 *Turdus* species (*T. pallidus*, brown-headed thrush [*T. chrysolaus*],
 217 eyebrowed thrush [*T. obscurus*], and Izu thrush [*T. celanops*]), but no deletion was detected in the
 218 other *Turdus* (Naumann's thrush [*T. naumanni*] and Japanese thrush [*T. cardis*]) and outgroup genera
 219 *Zoothera* (*Z. dauma*). Therefore, the East Eurasian thrushes can be divided into 3 groups according
 220 to the presence or absence of these synapomorphic indels. Eurasian *Turdus* Group 1 (ETG1) has
 221 characteristics of both tandem duplications and deletions, and includes *T. pallidus* and its conspecies
 222 *T. chrysolaus*, *T. obscurus*, and *T. celanops*. Eurasian *Turdus* Group 2 (ETG2) consists of the other
 223 major *Turdus* thrushes (*T. naumanni* and *T. cardis*) whose only tandem duplications are detected in
 224 the C-terminal domain. The thrush group who has neither tandem duplications nor the 3-amino acid
 225 deletion is considered to form an outer clade of ETG1 and ETG2 (Fig. 4) and named OT (other
 226 species of Turdidae).
 227 In the next step, we tried to identify to which group the other *Turdus* species belong to, according to
 228 the presence or absence of these synapomorphic indels. Two samples of the grey-backed thrush (*T.*
 229 *hortulorum*) collected in central Japan had the genotype of *RT2/RT2* and *RT2/RT3*, respectively, thus
 230 belonging to ETG2 (Table 2). The other 2 Eurasian *Turdus* species (island thrush [*T. poliocephalus*
 231 *niveiceps*] and rufous-throated thrush [*T. ruficollis*]) also had duplicated alleles that were
 232 characteristics of ETG2 thrushes, whereas the Siberian thrush (*T. sibiricus*) could be clearly
 233 distinguished from other *Turdus* species by the absence of a duplicated allele.

234

235

236 Discussion

237

238 We detected intraspecific polymorphisms in a variable number of tandem duplications in all *Turdus*
239 species (with large sample sizes), even though we did not detect a significant difference in the allelic
240 distribution between sexes ($P = 0.383$) nor between sampling locations (Honsyu vs. Kyusyu; $P =$
241 0.257) in *T. pallidus* populations. This simply suggests that tandem duplications in the C-terminal
242 tail would have no impact on the receptor functions, and that the current level of allelic distributions
243 is determined by a random genetic drift under the neutral mutation model of evolution. One of the
244 most plausible explanations for the retention of polymorphism within each *Turdus* species is gene
245 conversion events associated with genetic drift. Recent study suggests that gene conversion can
246 generate new haplotypes by transferring sections of DNA within and across duplicated loci in wild
247 bird populations (Spurgin et al. 2011). Similar mechanism may contribute to generate genetic
248 instability in the C-terminal domain of MTR. Gene conversion can also homogenize the level of
249 variation among populations by increasing or decreasing of the tandem duplication unit. At present,
250 however, we do not have an answer to the following fundamental questions: Why can only the
251 genomes of *Turdus* thrushes accommodate structural changes in the C-terminal domain of MTR?

252 Why are duplicated elements so strictly conserved among vertebrates? To address these questions,
253 we need to conduct site-directed mutagenesis and/or other biochemical experiments targeting
254 C-terminal domain of MTR.

255 The phylogenetic relationship inferred from synapomorphic indels strongly supports the conspecific
256 relationships among *T. pallidus*, *T. chrysolaus*, *T. obscurus*, and *T. celanops*. This is consistent with
257 molecular phylogenies based on the cytochrome c oxidase subunit I (*COI*) sequences (Saitoh et al.
258 manuscript in preparation) and datasets from 3 mitochondrial genes (Voelker et al. 2007). Moreover,
259 local ornithologists have reported that some of Eurasian *Turdus* belonging to EGT1 are temporally
260 intermingled with each other for wintering in Japan, and that *T. chrysolaus* sometimes can be seen in
261 the same flock with *T. pallidus*. All these findings suggest a close relationship and recent
262 diversification of EGT1 thrushes. In fact, captive and natural hybridization was reported between
263 several combinations of closely related *Turdus* species such as *T. chrysolaus* × *T. pallidus* (male) and
264 *T. naumanni* × *T. ruficollis* (McCarthy 2006, p. 239). Natural hybrid zones between Eurasian *Turdus*
265 species may help to maintain the allele frequency constant among each group of *Turdus* thrushes,
266 even though our data presented here clearly reject the possibility of hybridization between EGT1 and
267 EGT2 thrushes. *T. celanops* is an exceptional species, because it is considered a sedentary bird that
268 breeds and reproduces on small islands, in contrast to the other migratory thrushes. Therefore, *T.*
269 *celanops* is considered to have lost the rare alleles (*RT3*– and *RT4*–) recently due to local bottleneck

270 effects in some island populations.

271 Not only hybridization with closely related species but also other biological factors influence the

272 magnitude of polymorphisms in MTR. *T. pallidus* is a migratory bird of eastern Asia, and southern

273 and central parts of Japan are their primary wintering grounds (Brazil 1991). According to the first

274 breeding record in the Hiroshima prefecture (located in the western part of Honsyu) in 1991, some

275 populations of *T. pallidus* stay in Japan throughout the year (Ueno et al. 1993). Such a social and

276 behavioral dimorphism would promote diversification and speciation in local populations, resulting

277 in stochastic transitions of genetic variation. In our future research, the levels and patterns of

278 polymorphisms in other neurotransmitter-related genes will be examined between resident and

279 migrating populations of *T. pallidus*.

280

281

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295

296 Conflict of interest

297

298 The authors declare that they have no conflict of interest.

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373

Table 1

Distributions of wildtype and duplicated alleles among *Turdus* and *Zoothera* thrushes

Species/allele*	2n	NR	NR-	RT2	RT2-	RT3	RT3-	RT4	RT4-
<i>T. pallidus</i>	170		25 (0.15)		120 (0.71)		22 (0.13)		3 (0.02)
<i>T. chrysolaus</i>	22		6 (0.27)		15 (0.68)		1 (0.05)		—
<i>T. obscurus</i>	10		1 (0.1)		7 (0.7)		1 (0.1)		1 (0.1)
<i>T. celanops</i>	86		22 (0.26)		64 (0.74)		—		—
<i>T. cardis</i>	52	9 (0.17)		32 (0.62)		9 (0.17)		2 (0.04)	
<i>T. naumanni</i>	24	3 (0.13)		16 (0.67)		3 (0.03)		2 (0.08)	
<i>Zoothera dauma</i>	18	18 (1)		—		—		—	

374

375 * NR (wild-type, no duplication), RT2 (2 tandem repeats), RT3 (3 tandem repeats), and RT4 (4 tandem

376 repeats). A dash (—) beside allele name indicates the presence of the 3-amino acid deletion.

377

Table 2

Genotyping in *Turdus* species with less than 3 specimens

Species	<i>n</i>	Locations	Genotype(s)*	Group
<i>T. poliocephalus niveiceps</i>	1	Taiwan	<i>RT2/RT2</i>	ETG2
<i>T. ruficollis</i>	1	Mongolia	<i>NR/RT2</i>	ETG2
<i>T. hortulorum</i>	2	central Japan	<i>RT2/RT2</i> , <i>RT2/RT3</i>	ETG2
<i>T. sibiricus</i>	2	northern Japan	<i>NR/NR</i> , <i>NR/NR</i>	OT

378

379 * *NR* (wild-type, no duplication), *RT2* (2 tandem repeats) , and *RT3* (3 tandem repeats)

380

381 Figure Legends

382

383 **Fig. 1**

384 Schematic structure of the C-terminal tail of the mesotocin receptor (MTR). The double cysteine
385 residues for palmitoylation are highlighted and the amino acid residues highly conserved among
386 vertebrates are shown in capital letters. The position of the 3-amino acid deletion (Ser–Thr–Arg) is
387 boxed, and the tandem duplicated domain in *Turdus* thrushes is highlighted with gray background.

388

389 **Fig. 2**

390 Comparison of the amino acid sequences at the C-terminal tail of mammalian oxytocin receptors
391 (OXTRs), MTRs (amphibian, marsupial, and avian), and fish isotocin receptor. GenBank sequences
392 are derived from rat (rOXTR; NP037003.2), human (hOXTR; NP000907.2), cow (cOXTR;
393 NP776559.1), porcine (pOXTR; NP999192.1), canine (dOXTR; NP001185588.1), western clawed
394 frog (fMTR; XM002936297.1), gray short-tailed opossum (oMTR; XP001375059.1), green anole
395 (aMTR; XP003224939.1), non-passerine Neoaves (npMTR; AB634795—AB634816), zebra finch
396 (zfMTR; XM002188266.1), chicken (cMTR; NP001026740.1), and zebrafish (fISR;
397 XP001341507.1). RT2, RT3, and RT4 are duplicated elements found in *Turdus* thrushes. Highly
398 conserved motifs are shaded, and the symbols ¶ and § represent S/G/N/A/T and S/G, respectively.

399 The sequence of npMTR is truncated because of internal primer design.

400

401

402 **Fig. 3**

403 Alignment of amino acid sequences in the distal part of transmembrane VII and the C-terminal tail

404 of the avian mesotocin receptor. A dot or a dash denotes identity with the consensus sequence or a

405 deletion, respectively. The GenBank sequence of zebra finch (XM002188266) is also used for

406 alignment, and conserved residues in all avian taxa are highlighted. X in the sequence of Pthrush

407 [RT3–] indicates alanine or glycine due to heterozygous nucleotides in this position. A deletion that

408 could be identified only in Eurasian *Turdus* Group 1 (ETG1) is highlighted with gray background.

409 The square brackets indicate the number of repetitive units in the C-terminal tail, and duplicated

410 elements are shown at the bottom of the alignment (each repeat unit is boxed). Vertical bars represent

411 the following categories: P, passerine birds; N, non-passerine Neoaves; G, Galliformes; and A,

412 Anseriformes. Details of avian specimens are shown in S-Table 1.

413

414 **Fig. 4**

415 Phylogenetic relationships of East Eurasian thrushes inferred from synapomorphic characters at the

416 C-terminal domain of the mesotocin receptor. The timings of 3-amino acid deletion and 18-amino

417 acid duplications are indicated as red circle and boxes, respectively. Eurasian *Turdus* thrushes can be
418 divided into 2 groups, Eurasian *Turdus* Groups 1 and 2 (ETG1 and ETG2) according to these
419 synapomorphic characters.
420
421 Supplementary material
422 **Table S1** Details of avian specimens used for sequencing
423

		TM VII										C-terminal domain																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																																							
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Fig. 2

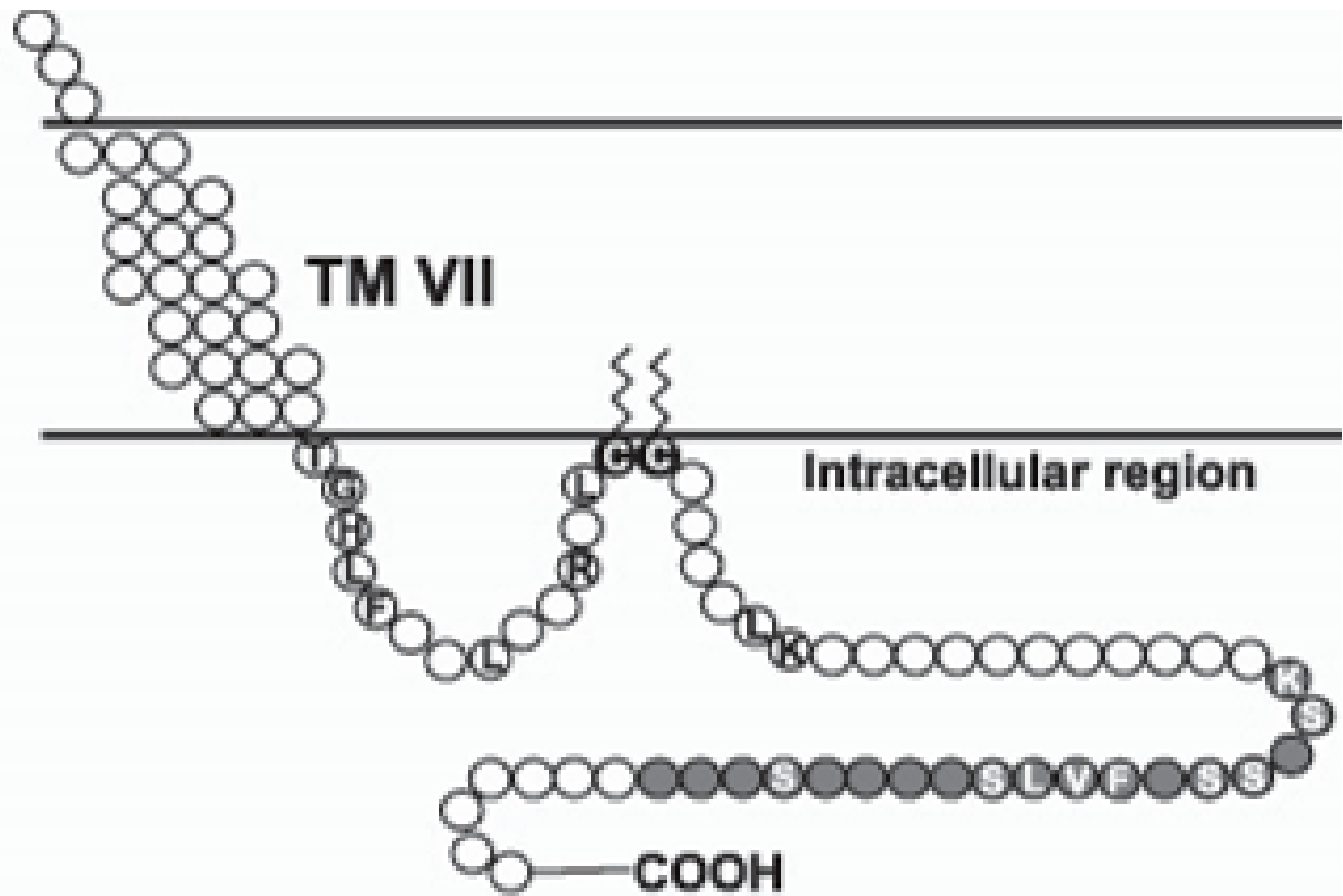


Fig.1

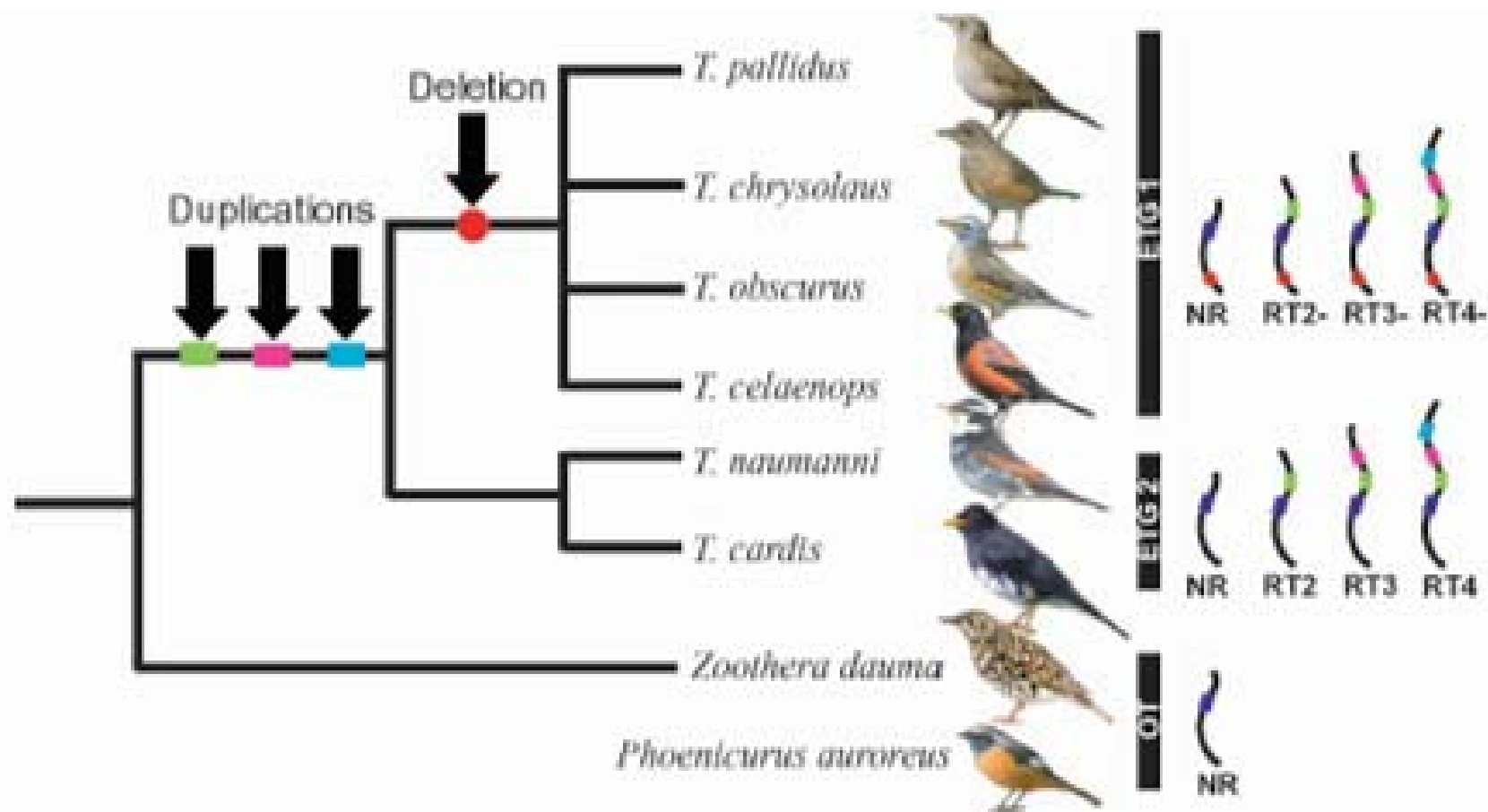


Fig. 4

Consensus	S	V	S	K	K	S	N	S	S	T	F	V	L	S	C	R	S	S	S	Q	R	S
rOXTR	•	•	•	•	•	•	•	•	•	•	•	•	•	•	R	•	•	•	•	•	•	•
hOXTR	•	A	•	•	•	•	•	•	•	S	•	•	•	•	H	•	•	•	•	•	•	•
cOXTR	•	•	•	•	•	•	•	•	•	•	•	•	•	•	Q	Y	•	•	•	•	•	R
pOXTR	•	•	•	•	•	•	•	•	•	•	•	•	•	•	Q	H	•	•	•	•	K	•
dOXTR	•	•	•	•	•	•	•	•	•	•	•	•	•	•	H	H	•	•	•	•	•	•
fMTR	•	T	•	R	•	•	•	•	•	•	•	•	•	•	R	K	•	•	•	•	K	•
aMTR	•	•	•	•	•	•	•	•	•	•	•	•	•	•	L	K	•	•	•	•	•	•
oMTR	•	A	•	•	•	•	•	•	•	•	•	•	•	•	R	K	•	•	•	L	K	•
npMTR	¶	•	§	•	•	•	•	•	•	S	•	•	•	•	•	K	[reverse primer]					
zfMTR	–	–	•	•	•	•	•	•	•	S	•	•	•	•	•	•	G	T	•	H	•	•
RT2/3/4	–	–	F	Q	•	•	•	•	•	S	•	•	•	•	•	•	•	P	•	H	•	•
cMTR	•	•	G	R	•	•	H	•	•	S	•	•	•	•	•	•	•	•	•	•	•	•
fISR	Q	D	•	R	•	•	•	•	•	T	Y	•	I	–	K	S	T	•	•	•	•	•

Fig. 3